Expression of Heat Shock Protein (HSP A1A) and MnSOD Genes Following Vitrification of Mouse MII Oocytes with Cryotop Method

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Abstract

Objective: The aim of this study was to investigate the effects of two vitrification protocols on mouse metaphase stage II (MII) oocytes and evaluate their effects on the expression of heat shock protein A1A (HSP A1A) and MnSOD genes.

Materials and Methods: Groups of approximately 15 oocytes without cumulus complexes were collected and vitrified with 10% (v/v) ethylene glycol (EG) + 10% (v/v) dimethyl-sulphoxide (DMSO) + 0.5M sucrose in group A (VSI) and 14.5% (v/v) EG + 14.5% PROH + 0.5M sucrose in group B (VSII), respectively. Thawing after vitrification was performed by placing the oocytes into 1M sucrose for 1 minute and two diluted solutions, each for 3 minutes. After thawing, the oocytes were fertilized and cultured in vitro to develop into the pronuclear stage. The survival rate of vitrified-warmed oocytes and rate of fertilization were evaluated. In addition, gene expressions (HSP A1A, MnSOD and β-actin) of vitrified-warmed oocytes were also examined by reverse transcription polymerase chain reaction (RT-PCR).

Results: Survival rates of each group were separately compared to the control. The result showed significant differences between each experimental group compared to the control (p ≤ 0.001). The survival rate of oocytes after warming was higher in VSI (91.2% ± 1.7) compared to VSII (89.2% ± 1.5) but there were no significant differences between the two groups. The rate of fertilization was significantly (p ≤ 0.05) reduced in vitrified-warmed (VSI: 39% ± 5.8; VSII: 34% ± 5.7) oocytes compared to the control (88.36% ± 2.3). The expression of MnSOD increased in the vitrified-warmed oocytes when compared to control oocytes. We also detected HSP A1A only in the control and VSI group.

Conclusion: Vitrification of oocytes by cryotop resulted in high survival rate; low developmental competence and fertilization rate of vitrified-warmed oocytes which may be a result of changing expression of important genes after thawing.

Keywords: Oocyte, Vitrification, Cryotop, Gene Expression, HSP A1A, MnSOD

Introduction

Oocyte cryopreservation as an essential part of assisted reproductive technology (ART) has recently received major success in clinical practice (1, 2). This technology is used to preserve fertility in young women that are at risk of losing ovarian function for a variety of reasons such as chemotherapy or radiotherapy for cancer and autoimmune or hematologic diseases (3, 4).

Although sperm and embryo freezing has greatly improved, oocyte cryopreservation still faces numerous problems (5). At present, vitrification is a feasible method for cryopreservation of oocytes or embryos (2). In this procedure, glasslike solidification is achieved by using a high concentration of cryoprotectants (CAPs) and a rapid cooling rate (6). For the first time, Rall and Fahy used vitrification to freeze mice embryos (7). Several vitrification protocol were proved according to different types of tools, including the open pulled straw (8), electron microscope grid (9), cryoloop (10), and cryotop (11) in addition to different cryoprotectant solutions (6, 12).

Among the various cryodevices, the cryotop as devised by Kuwayama has recently attracted more attention since a minimum volume of solution (ap-
proximately 1μl) is loaded onto it, thus achieving both a high cooling (-23000°C/min) and warming (40000°C/min) rate(15). Furthermore the concentration of permeable CPAs is decreased to 30% by this method, thus minimizing the potential toxic effect (13-15). Based on further studies the cryotop is an easy and reliable procedure (15). Lucena et al.(16) achieved an 89% survival rate and up to 56% pregnancy rates with this method.

The most common CAPs used for the vitrification method are ethylene glycol (EG), dimethylsulphoxide (DMSO) and 1,2-propanediol (PROH) as permeable and sucrose as nonpermeable CAPs (17, 18). Kuwayama et al. have reported that the combination of DMSO and EG in equal proportions was the most efficient (15). Furthermore, the low molecular weight, high permeation ability and low toxicity of EG have enabled it to be used for vitrification of human oocytes and embryos. DMSO can facilitate EG permeability, cause increased polyplody and may cause spindle polymerization whereas PROH is more permeable compared to DMSO (19). In addition, Vajta et al. have reported that cryoprotectant mixtures when compared to solutions that contain only one permeable cryoprotectant can decrease individual specific toxicity (8, 20). Moreover, osmotic stress may be induced by the use of a high concentration of CPAs (21), thus, the combination of a low concentration of CAPs can reduce this negative effect.

During the past decade, further studies have reported only structural and morphological damages such as zona hardening, modification in selective permeability of the plasma membrane, aneuploidy and nuclear fragmentation in the vitrified-warmed oocytes during vitrification (5).

Accordingly, there is little information on biological and molecular events following cryopreservation of oocytes (5, 22). Heat shock protein (HSP A1A has a protective function by heat or stress response on the cellular auto regulation. The critical role of HSP A1A as a protective protein due to external stress has been proven. It has been demonstrated that knock out HSP 70.1 mice had higher sensitivity to osmotic stress after preconditioning with heat was decreased (23-25). Hut et al. observed that HSP A1A has a protective role in the mitotic cell and prevented chromosomal division that occurs following heat-induced centrosome damage (26). Manganese super oxide dismutase (MnSOD) is an anti-oxidant enzyme that protects oocytes and embryos against oxidative stress damage. It has been shown that addition of antioxidant enzymes such as catalase or SOD1 (Cu-Zn-SOD) to culture media lead to an increased rate of blasto-cyst formation in the rabbit (27), mouse (28), and bovine species (29).

Sona et al. have reported that cold stress can be influenced on expression of several genes that are related to stress (30). Succu et al. (31) found that vitrified oocytes have shown low developmental competence after in vitro fertilization (IVF) and in vitro culture (IVC). Thereby for evaluating the post warming quality of oocytes, this is the best way to focus on alternation of gene expression related to stress and important cell function in response to cooling.

To the best of our knowledge, gene expressions have not been completely investigated in mammalian vitrified oocytes. The objective of present study was to determine expression of some genes related to stress including HSP A1A and antioxidant enzyme MnSOD. On the other hand, the efficiency of the cryotop method with the mixture of CAPs on survival, morphology, IVF and gene expression of mouse oocytes were examined.

Materials and Methods
All chemicals and media were purchased from Sigma-Aldrich Co. (St. Louis, Mo, USA), unless otherwise indicated.

Oocyte collection
In this study, 8-10 week old female mice (NMRI strain) were used. The mice were kept under the a 12 hours light: 12 hours dark cycle for two weeks to adapt to laboratory conditions. The animals were superovulated by intraperitoneal injections of 10 IU PMSG and followed 48 later by 10 IU HCG. The mice were sacrificed by cervical dislocation 13-15 hours post-HCG administration.

The oocyte cumulus complexes were taken from the ampulla of oviduct and put in hepes-buffered M199 medium + 20% FBS. Denudation of oocytes were done by placing the oocyte cumulus complexes in M199 supplemented with 20% FBS was used as a carrier for the cryoprotectants. Denuded MII oocytes were vitrified in groups of 15 by the minimum volume cooling (MVC) method using the cryotop as a device (14). The complete vitrification process was accomplished at room temperature (25°C). In the present study, a mixture of two concentrations
of vitrification solution (VS) were used. In group A, VS1 supplemented with 10% ethylene glycol (EG) + 10% dimethylsulphoxide (DMSO) + 0.5 M sucrose in base medium. In group B, VS2 consisted of 14.5% EG + 14.5% PROH in base medium. The equilibration solutions were 50% VS1 and VS2 which contained no sucrose. Oocytes were primarily exposed to first equilibration drop for 3 minutes, after which they were merged with an adjacent ES drop. Subsequently, the oocytes were transferred to the vitrification solution for 1 minute. Then, oocytes were quickly loaded onto the top of the cryotop (Kitzato Ltd. Tokyo, Japan). All excess media were carefully removed and the cryotops were promptly submerged in liquid nitrogen (LN2).

**Thawing**

Upon warming, the cryotops were instantly inserted into the thawing solution (1M sucrose, M199 + 20% FBS) for 1 minute at 37°C. Subsequently, oocytes were placed into decreasing sucrose concentrations (0.5 M, 0.25 M) for 3 minutes each, then twice washed in holding medium for 5 minutes for cryoprotectant removal. The survival rate of vitrified-warmed oocytes was assessed on the basis of normal appearing zona pellucida, an integral plasma membrane and intact polar body as visualized under light microscope.

**IVF**

Survived epididymal vitrified, warmed oocytes were chosen and placed into 200 μl drops of IVF medium (HTF + 15 mg/ml BSA) layered with mineral oil. The medium was prepared previously and equilibrated at 37°C in a 5% CO₂ incubator. Suspensions Ham's F-10 of epidydimal sperm were incubated for 45-60 minutes in capacitation medium (ham's F10 + 4 mg/ml BSA). A final concentration of 2 × 10⁶ spermatozoa/ml were added to IVF medium which contained 10-15 oocytes and incubated at 37°C in 5% CO₂. After 26 and 32 hours post-fertilization, the number of cleaved oocytes that developed into the pronuclear stage was recorded.

**RNA extraction**

Total RNA was extracted from 20 vitrified and non-vitrified oocytes with tripure isolational reagent (Roche, Germany) according the manufacturer’s instructions. The concentration of extracted RNA and purity were determined by ND-1000 spectrophotometer (Nonodrop, Wilmington, DE). RNA was subsequently subjected to reverse transcription polymerase chain reaction (RT-PCR) which was performed using cycle script RT premix (dN) reverse transcriptase (Bioneer, South Korea) with 300ng of total RNA following the manufacturer's protocols.

**PCR**

The cDNA equivalents of 20 oocytes were used in each reaction. PCR was performed using Taq polymerase enzyme (Roche, Germany). PCR analysis was carried out in a total volume of 25μl that consist of 1μl of each primer mix, 2μl dNTP, 2.5μl 10x buffer with MgCl₂, 0.3μl rTaq polymerase enzyme, 1μl cDNA and 18.2μl DWW. After an initial denaturation step of 3 minutes at 94°C, amplification of 20 oocytes was performed. Denaturation in each cycle included: 94°C for 30 seconds and annealing at 55 °C for MnSOD, and 59°C for HSP A1A for 30 seconds and extension at 72°C for 1 minute.

RT-PCR products underwent electrophoresis on a 2% agarose gel, then were stained by ethidium bromide (Cina gene, Iran) and visualized under ultraviolet.

**Statistical analysis**

Oocyte survival and fertilization rates were analysed by ANOVA. Vitrification and *in vitro* fertilization data were analyzed by SPSS version 11.5 software.

**Results**

**Vitrification and IVF**

The results are summarized in table 2. After thawing, the survival rate of oocytes was high and there was no significant difference between VSI (91.2% ± 1.7) and VSII (89.2% ± 1.5). Survival rate of each group was seperately compared to the control. The results showed significant differences between each experimental group compared to the control (p ≤ 0.001). The survival rate of oocytes after warming was higher in VSI (91.2% ± 1.7) compared to VSII (89.2% ± 1.5) but there were no significant differences between the two groups. The survival rates of oocytes were higher in VSI (91.2% ± 1.7).

As shown in table 2, the *in vitro* fertilization rates were significantly decreased in vitrified-warmed oocytes (VSI: 39% ± 5.8; VSII: 34% ± 5.7) compared to the control (88% ± 2.3). Although the fertilization rates were not different between the
two experimental groups, the results showed significantly lower fertilization rates compared to the control ($p \leq 0.05$). The fertilization rate was higher in VSI (39\% ± 5.8) compared to the VSII (34\% ± 5.7) group.

**Gene expression analysis**

As shown in figure 1 the expressions of all genes in the vitrified-warmed oocytes were compared to the control. RT-PCR was prepared to investigate the alternation in gene expressions. The abundance of mRNA was generally reduced in oocytes related to vitrification procedures, but expression of Mn-SOD increased in vitrified-warmed oocytes when compared to control oocytes. HSP A1A was only detected in the control and VSI group.

**Discussion**

In the present study we observed that the expression of a novel member of HSP70 family, HSP1A1, and an antioxidant enzyme, MnSOD, were altered in vitrified-warmed MII oocytes when compared to the control. A decrease in fertilization rate of the vitrified-warmed group after IVF was also seen. Intrinsic factors in oocytes are responsible for controlling the rate and time of cleavage. According to studies that have been performed in the animal model, decreasing fertilization rate and low developmental competence of oocytes after warming may be associated with alternations in expressions of antioxidant enzymes (32, 33) and cryoprotectant toxicity (34, 35). The developmental competence of oocytes depend on the abundance of specific gene transcripts (36-38).

The genes we chose are involved in response to stress (MnSOD, HSP A1A) and constitutive function of the cell ($\beta$-actin). Changes in gene expression are considered as an integral part of the cellular response to thermal stress. It is widely accepted that HSPs are the best candidates whose expressions are affected by heat shock, moreover it has been recently shown that thermal stress also induced expression of a number of non-HSP genes such as MnSOD (30). Briefly, heat stress impacts different cell functions such as: 1) inhibition of DNA synthesis, transcription, RNA processing and translation; 2) disruption of cytoskeletal component; and 3) changes in membrane permeability that cause an increase in Ca$^{2+}$, Na$^+$ and H$^+$ (39, 40).

HSP A1A, a member of the inducible heat-shock family, can protect oocytes against different drastic conditions, including oxidative stress (41, 42). In the current study, we demonstrated that the expression of HSP A1A reduced in vitrified-warmed MII when compared to control oocytes; we detected HSP A1A only in the control and VSI groups. Boonkusol et al. also reported similar results after vitrification with straw. This might be a sign of decreasing oocyte viability following vitrification.
was obtained in the present study confirmed the expression of specific genes (3). The result that process may alter DNA integrity and affect the cellular function and affect further developmental competence of oocytes (46). Oxidative stress can cause DNA instability in mouse oocytes. Moreover, Bilodeau et al. have reported that during the cryopreservation the activity of SOD was reduced to 50% in bovine spermatozoa (47). Therefore high expression of MnSOD in vitrified-warmed oocytes can be a defense mechanism against oxidative stress (46). All vitrification processes may be induced by stress. There are several critical factors that affect vitrification such as the concentration and toxicity of cryoprotectants, protocol and cryodevice. Therefore it is important to choose a proper approach in order to minimize oxidative, osmotic and heat stress (48). For the first time, Kuwayama et al. designed the cryotop and reported a 91% survival rate, 81% cleavage rate and 50% blastocyste rate result to 11 live births in humans (14). According to further studies that have been recently completed, the efficiency of the cryotop technique for cryopreservation of sensitive samples, such as human oocytes was confirmed. Additionally, high survival and blastocyste production rates were achieved (20, 49).

In the current study, an attempt was made to increase the cooling rate by using the minimum volume cooling method (cryotop). Additionally we attempted to decrease cytotoxicity of the cryoprotectant with the vitrification solution and therefore reduce stress. For the first time, Ishimori et al. successfully used a combination of EG and DMSO. In order to prevent intracellular ice crystal formation, permeating cryoprotectants are used. On the other hand, using a combination of two or three CAPs are effective to reduce the concentration and individual specific toxicity of each CAP (20).

Besides, osmotic stress during the vitrification process may alter DNA integrity and affect the expression of specific genes (3). The result that was obtained in the present study confirmed the efficiency of the cryotop method in vitrification of MI oocytes. It has been recently reported that using a mixture of 15% EG and 15% DMSO is beneficial for vitrifying human oocytes. Chian et al. also demonstrated the efficiency of a mixture of 15% EG and 15% PROH (4). During the Vitrification procedure the equilibration time, thawing and concentration of cryoprotectant vary in relation to both the species and laboratory condition. Therefore, in the current study we compared two mixtures of 14.5% EG + 14.5% PROH and 10% EG + 10% DMSO in mouse MII oocytes. In contrast to the studies that have been undertaken by Chian et al. we found that the survival rate and developmental competence of mouse MII oocytes improved after vitrification in the mixture of 10% EG + 10% DMSO when compared to 14.5% EG + 14.5% PROH. These results support the hypothesis that the concentration of each cryoprotectant was reduced when mixed and thus can be effective in decreasing stress (30).

Conclusion
In conclusion, our finding in the present study showed the potential of the cryotop in vitrification of mouse MII oocytes. A possible reason for low developmental competence and cleavage may be the low abundance of mRNA after vitrification. In any case, it is essential to perform further studies to focus on expression of genes that are involved in the crucial practice of cells in different stages.

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References
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